Lactococcus lactis NRRL B-30656 and the biopolymer production process

Background of the invention

1. Field of the invention

This invention relates to a glucose and fructose polymer and the method for preparing it using a *Lactococcus lactis* strain. The exopolysaccharides are natural glucose and fructose polymers. These polymers can be found in several plants and microorganisms and are useful as emulsifiers, thickener and surfactants in the food and medicaments industries.

2. Description of the state of the art

Fructosans naturally occur in two general forms differentiated by the type of binding between molecules of fructose: inulin, as found in plants, is formed from a column of fructose molecules bound by beta,2-1 links. Levans,formed as microbial products, have a column of fructose molecules bound by beta,2-6 links. The fructosans from plants are smaller (around 100 residues) whilst microbial levans contain more than 3 million residues (Pontis et al., 1985, Biochemistry of Storage Carbohydrates in Green Plants. In: Dey and Dixon (eds). Ch. 5, p. 205. New York, Academic Press).

Microbial Levans are produced with sucrose-based substrates having a variety of microorganisms: Acetobacters (Loewenberg, et al., 1957. Can. J. Microbiol., Vol. 3, p. 643), Achromobacter sp. (Lindberg, G., 1957. Nature. Vol. 180, p. 1141),

Aerobacter aerogenes (Srinivasan, et al., 1958. Science. Vol. 127, p. 143),
Phytobacterium vitrosum (Belval, et al., 1947. 1948. Compt. Rend. Vol. 224, p. 847
and Vol. 226, p. 1859), Xanthomonas pruni (Cooper, et al., 1935. Biochem. J. Vol. 29, p. 2267), Bacillus subtilis (Dedonder, R., 1966. Meth. Enzymol. Vol. 8, p. 500 and
Tanka, et al., 1979. J. Biochem., Vol. 85, p. 287), Bacillus polymyxa (Hestrin et al., 1943. Biochem. J., Vol. 3, p. 450), Aerobacter levanicum (Hestrin, et al., Ibid.),
Streptococcus sp. (Corrigen et al., 1979. Infect. Immun., Vol. 26, p. 387),
Pseudomonas sp. (Fuchs, A., 1956. Nature. Vol. 178, p. 92) and Corynebacterium
laevaniformans (Dias et al., 1962. Antonie Van Leewenhoeck, Vol. 28, p. 63).

There are some reports of levan being produced at very low levels and having low purity to be used industrially.

Other biological polymers such as xantan and dextran gum have been extensively used in the food industry as stabilisers in emulsions and froth in ice-cream, in salad-dressing, etc. (Sharma, S.C., January 1981. J. Food Tech., p. 59). Extracellular polysaccharides produced by microorganisms offer a variety of uses and potentially low costs.

Small quantities of levan are generally produced by sucrose fermentation using Actinomyces viscosus or Aerobacter levanicum strains.

Bacillus polymixa generally produces hetero-polysaccharides having different forms of polymers. Genetically modified E. coli strains have been used for producing levan (Gay, P. et al., 1983. J. Bacteriol. Vol. 153, p. 1424). Furthermore, other aerobic

fermentation methods have also been used for producing levan (Jeanes, et al., U.S. Pat. No. 2,673,828; Gaffor, et al., U.S. Pat. No. 3,879,545; Ayerbe, et al., U.S. Pat. No. 4,399,221). The drawback of such processes is that they produce low product yield and problems related to contamination, thereby industrial processes leading to greater productivity are required.

Description of the invention

The main purpose of this invention was to provide a biopolymer produced by an enzyme transferase having glucose and fructose transfer activity. It was produced from a Lactococcus lactic strain (NRRL B 30656) characterised by its high transfer activity, allowing the obtention of the biopolymer by a simple production method which was easy to scale-up. Its method of production consisted of the following steps: Phase 1: fermentation with the Lactococcus lactis NRRL B 30656 strain in culture medium developed for this micro organism's growth. Phase 2: extracellular enzyme recovery by centrifuging or ultra-filtration. Phase 3: biopolymer production by enzyme reaction using sucrose as substrate and enzyme extract. Phase 4: biopolymer purification by precipitation with solvents or ultra-filtration followed by drying the product.

The main purpose of this invention is to provide a biopolymer, produced by an enzymatic extract or preparation having glucosyltransferase and fructosyltransferase activity. It is produced from a Lactococcus lactis strain (NRRL B-30656) characterised by its high transfer activity, allowing the biopolymer to be obtained by a simple production method which is easy to scale-up. Its production comprises the following steps: Phase 1: fermentation with the Lactococcus lactis NRRL B-30656

strain in a culture medium developed for this microorganism's growth; Phase 2: extracellular enzyme recovery trough centrifuging or ultra-filtration; Phase 3: biopolymer production trough enzyme reaction using sucrose as substrate and the enzymatic extract or preparation; and Phase 4: biopolymer purification trough precipitation with solvents or ultra-filtration followed by drying the product.

Detailed description of the invention

The object of the invention is to produce a polysaccharide contaminants-free pure biopolymer. The biopolymer can be described as being a polymer produced by a Lactococcus lactis strain isolated from soil. This strain has high transfer activity, leading to obtaining the biopolymer through a simple process, having a purity greater than 95%. The object of the invention was to produce a pure biopolymer which was free from polysaccharide contaminants. The biopolymer can be described as a polymer produced by a Lactococcus lactis strain isolated from the soil. This strain has high transfer activity, leading to obtaining the biopolymer by a simple process, having greater than 95% purity.

The microorganism. The Lactococcus lactis NRRL B-30656 strain is isolated from soil in the present invention by a selective process using a sucrose-containing medium sucrose-as a carbon source in which the microorganisms producing the enzymatic extract or preparation having glucosyltransferase and fructosyltransferase activity, are able to use the substrate and to produce the polymers, giving the colony a mucoid aspect. Microorganisms having these characteristics are selected from this medium and purified trough isolating techniques involving successive dilutions and

The microorganism. The Lactococcus lactis NRRL B-30656 strain was isolated from soil in the present invention by a selection process using a medium containing sucrose as earbon source in which transferase enzyme producing microorganisms were able to use the substrate and produce polymers, giving a mucoid aspect to the colony. Microorganisms having these characteristics were selected from this medium and purified by isolation techniques involving successive dilutions and plate isolation. The Lactococcus lactis NRRL B-30656 strain was obtained from these strains and was used in the present invention.

In accordance with the present invention, the Lactococcus lactis NRRL B-30656 strain has been deposited in the Agricultural Research Service Patent Culture Collection NRRL Reference Bank; it was assigned registration number NRRL B-30656 by this institution. This strain produces an enzyme having 2-6 U/ml glucose transfer activity, using sucrose as substrate and also produced a 900-1,100 K Dalton molecular weight glucose and fructose polymer.

The strain <u>iswae-called NRRL B-30656</u>. This strain was isolated and characterised at the Universidad Nacional de Colombia's Instituto de Biotecnología (IBUN). The strain <u>iswae-kept</u> at 4°C in Petri dishes with a culture medium whose composition <u>iswae</u>: 10-40 g/l sucrose, 7-30 g/l yeast extract, 5-20 g/l 0.05-05 g/l potassium phosphate, 10-100 ppm mineral salts, pH 5-9.

The microorganism was characterised by optical microscopy using Gram staining and electronic transmission microscopy by means of positive staining with uranyl acetate and lead citrate. The biochemical characterisation was done using the computerised MicroScan system, according to that described in Bergey's determinative bacteriology

manual (Stanley, W; Sharpe, E; Holt, J. 1994. Bergey's Manual of Systematic Bacteriology, William and Wilkins, Baltimore).

Culture medium. A balance was carried out between carbon source, nitrogen source and certain trace elements for designing and optimising the culture medium for the fermentation with the NRRL B-30656 *Lactococcus lactis* strain. The culture medium provideged the microorganism with the nutrients needed for growing and producing the enzyme.

The following concentrations were established, as a result of evaluating culture medium components:

Component	Concentration (g/l)	
Salts		
K ₂ HPO ₄	7-30	
FeSO ₄ . 7H ₂ O	0.01-1	
MgSO ₄ . 7H ₂ O	0.01-0.1	
MnSO ₄ . H ₂ O	0.001 - 0.1	
CaCl ₂ . 2 H ₂ O	0.001 - 0.01	
NaCl	0.01-0.1	
Carbon source		
Sucrose	10-40	
Nitrogen source		
Yeast extract	7-30	

The pH is set to pH 5-9 with HCl. The medium is sterilised at 121°C for 15
minutes. The pH was adjusted to pH 5-9 using HCl. The medium was sterilised at
121°C for 15-minutes.

Fermentation. The pre-inoculums corresponding to 5-20% of the inoculum volume are activated from the pure strain preserve at -70°C in a medium having 20% glycerol; incubation time should not exceed 10-36 hours during which time pre-inoculum purity must be verified. These cultures are done in flasks with stirring, occupying 5-20% total volume; they are incubated at 20-40°C with 100-400 rpm stirring rate in orbital agitators. The number of inoculums necessary is determined by the number and size of the fermenters. Fermentation. The pre-inoculums corresponding to 5-20% inoculum volume were activated from the pure-strain kept at -70°C in the medium with 20% glycerol; incubation time did not exceed 10-36 hours, during which time it was necessary to verify the pre-inoculum purity. These cultures were done in stirring flasks, occupying 5-20% of the total volume and incubating them at 20-40°C with 100-400 rpm shaking in orbital shakers. The number of inoculums necessary was determined according to the number and size of the fermenters.

Growth and enzyme production conditions are 20-40°C temperature with stirring at a rate of 100-400 rpm (depending on the fermentation scale). Growth-and-enzyme production conditions were: 20-40°C temperature and 100-400 rpm-agitation (depending on the fermentation scale).

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Aeration. The fermentation promoting microorganism is aerobic, meaning that the

culture had to be aerated with 0.1 -1 volumes of air per medium volume per minute

(vvm) and pH is kept between 5 and 9 during fermentation. Culture mediums

resulting from this production process have combinations of components in order to

obtain final biomass concentration of 10-30 g/l, a wet weight, having 2-6 U/ml

transfer activity, this being achieved in 6-24 hours. Aeration. The microorganism that

enhances the fermentation is aerobic, meaning that the culture had to be aerated with

0.1 1 volumes of air per-medium volume per minute (vvm) and pH kept between 5

and 9 during fermentation. Culture mediums resulting from this production process

had combinations of components for reaching final 10 30 g/l biomass concentration,

wet weight, having 2-6 U/ml transfer activity, this being achieved in 6-24 hours.

Enzyme recovery. Extra-cellular enzyme are collected from fermented culture

medium through centrifuging at 3,000 -10,000 rpm for 15 minutes or by separating

the biomass through filtration. Enzymatic extract or preparation thus presents 2-6

<u>U/ml glucosyltransferase and fructosyltransferase activity.</u> Enzyme recovery.

Extracellular enzyme was recovered by centrifugating at 3,000 -10,000 rpm for 15

minutes or filtration to separate the biomass. Enzyme extract thus presented a 2-6

U/ml transferase activity.

Biopolymer production

Enzymatic reaction. Reaction conditions wereare-as follows:

Reaction medium:

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50-300 Mm phosphate buffer pH : 5-9

Substrate : 5-40% sucrosea

Ouantity of enzyme : 10-40% v/v enzymatic extract or

preparation

Reaction time : 12-48 hours

StirringAgitation : 100-400 rpm

Enzyme reaction. Reaction conditions were as follows:

Reaction medium:

50-300 Mm phosphate buffer pH : 5-9

Substrate : 5-40% sucrose

Quantity of enzyme : 10-40% v/v of enzyme extract

Reaction time : 12 48 hours

stirring : 100-400 rpm

Biopolymer recovery and purification

After the enzymatic reaction, the temperature was reduced to 4°C following enzymatic reaction and the biopolymer was recovered in two ways: The temperature was reduced to 4°C following enzymatic reaction and the biopolymer could be recovered in two ways:

a) Precipitation with solvents

96% ethanol iswas added to the cold reaction mixture with -stirringagitation. The added amount of ethanol corresponds to 1.2-2.0 volumes of ethanol/ reaction mixture

volume,96% ethanol was added to the cold reaction mixture with stirring. The quantity of ethanol added was 1.2 - 2.0 volumes of ethanol/reaction mixture volume.

The precipitated biopolymer <u>is</u>was redissolved in half the volume of deionised and distilled water and precipitated again with 1.2-2.0 volumes of ethanol/ reaction mixture volume.

The precipitated biopolymer <u>is</u>was redissolved in a third the volume of water and dried by lyophilisation or dried by compressed air at 60°C until reaching 5-6% Humidity.

b) Ultrafiltration

The reaction mixture <u>is</u>was submitted to ultra-filtration through a regenerated cellulose membrane having a pore size greater than 10,000 Dalton to eliminate residual glucose and fructose. The biopolymer <u>is</u>was then submitted to aspersion drying.

The biopolymer <u>is</u>was characterised by high performance liquid chromatography and 10% solution viscosity at 30°C. The biopolymer present<u>sed</u> a 7-7.5 minutes retention time using a Shodex SC1011 column at 70°C, 0.6 ml/min flow and HPLC grade water as mobile phase.

The viscosity of a 10% solution at 30oC was found to range from 1000-3000 centipoises (cP) using a ViscoEasy viscosimeter (Serie L, Schott, Ref. 28.541.120) L2 stem at 50 rpm. The viscosity of a 10% solution at 30°C was found to range from 400-

800 centipoises (cP) using a ViscoEasy viscosimeter (Serie L, Schott, Ref. 28.541.1201 L2 stem at 50 rpm.

Average DVS (diameter/volume/surface) particle size <u>is</u>was 224 micron. The biopolymer hased a true density close to that of ssucrose (1.5 mg/ml). It is a material presenting high inter-particle porosity (48%).

Examples

The following examples are given to illustrate the present invention.

Example 1

Isolating and identifying the biopolymer-producing microorganism

A biopolymer-producing bacterium was isolated from soil and identified as being Lactococcus lactis NRRL B-30656. 10 g samples were collected from soil and grown in 100 ml liquid medium containing sucrose as carbon source. This was incubated at 30°C with stirring for 24 hours. 4 x 1:10 dilutions were done in saline solution once growth was obtained; the fourth dilution was seeded. This culture was re-seeded in solid medium using the same composition and isolations were done, selecting the colonies showing polymer production. The culture was then transferred to a fresh medium and cultured for 24 hours. The microorganism was kept in a sucrose medium with 20% glycerol at -70°C and by lyophilisation using 10% skimmed-milk, once it had been isolated.

The isolated strain, cultivated in solid sucrose medium, showed the following macroscopic characteristics: clear, cream-coloured, rubbery, circular colonies having a defined edge of around 2 to 3 mm diameter (in 24 hours culture). Gram cocci were observed by microscope via Gram staining; they were occasionally found individually but were generally seen forming groups.

Electronic transmission microscopy characterisation led to observing circular cells in which the cell wall could be differentiated. No special structures were observed (i.e. electro-dense granules, flagella, fimbria, etc).

The strain of the present invention is *Lactococcus lactis* NRRL B-30656, catalogued as GRASS microorganism and shows the following biochemical characteristics:

Test	Result
Growth at 10°C	Positive
Growth at 15°C	Positive
Growth at 42°C	Negative
Growth at pH 4.8	Positive
Growth at pH 6.5	Positive
Growth at pH 9.2	Doubtful
Growth in 0.5% NaCl	Positive

Growth in 4% NaCl	Positive	
Growth in 5% NaCl	Positive	
Growth in NaCl 6.5%	Positive	
Growth in 10% NaCl	Negative	
Growth in 15% NaCl	Negative	
Catalase	Negative	
Haemolysis	Gamma	
Motility	Negative	
Vogees-Proskauer	Positive	
Aerobic glucose	Positive	
Anaerobic glucose	Positive	
Gas production	Negative	
Aerobic lactose	Positive	
Anaerobic lactose	Positive	
Gas production	Negative	
Aerobic fructose	Positive	
Anaerobic fructose	Positive	
Gas production	Negative	
Aerobic maltose	Positive	
Anaerobic maltose	Positive	
Gas production	Negative	

Aerobic manitol	Doubtful		
Anaerobic manitol	Doubtful		
Gas production	Negative		
Aerobic galactose	Positive		
Anaerobic galactose	Positive		
Gas production	Negative		
Aerobic sucrose	Positive		
Anaerobic sucrose	Positive		
Gas production	Negative		
Aerobic xylose	Doubtful		
Anaerobic xylose	Doubtful		
Gas production	Negative		
Aerobic rafinose	Positive		
Anaerobic rafinose	Positive		
Gas production	Negative		
Ribose	Positive		
Trealose	Positive		
Sorbitol	Positive		
Mannose	Positive		
Arabinose	Positive		
Arginin	Positive		

Example 2

Extract production or enzymatic preparation

1. Fermentation:

a) Microorganism activation

The Lactoccoccus lactis NRRL B-30656 microorganism was used for obtaining an enzymatic extract or preparation having glucosyltransferase and fructosyltransferase activity. Bacteria were stored in a cryoprotection solution (glycerol) at -70°C. The strain was slowly unfrozen until room temperature was reached and it was activated in 50 ml sucrose medium at 30°C for 12 hours and stirring at 180 rpm. 5 ml of this culture were used for two types of seeding. The first in agar sucrose suechrose, incubated at 30°C for 24 hours, while observing its mucoid characteristics and then stored at 4°C; the second in 100 ml sucrose broth incubated at 30°C for 12 hours. The latter was distributed in 1 ml centrifuge tubes with 20% v/v glycerol and stored at -70°C, for later use in fermentations. The remaining 45 ml of initial culture were preserved in 5 ml vials, lyophilised using 10% concentration sterile skimmed milk as support and stored at 4°C.

The Lactoccoccus lactis NRRL B 30656 microorganism was used for obtaining the transferase enzyme. Bacteria were stored in a eryoprotection solution (glycerol) at 70°C. The strain was slowly unfrozen at room temperature and activated in 50 ml sucrose medium at 30°C for 12 hours and stirring at 180 rpm. 5 ml of this culture were used for two types of seeding. One was seeded in sucrose agar, incubated at 30°C for 24 hours, mucoid characteristics were observed and stored at 4°C; the second was seeded in 100 ml sucrose broth and incubated at 30°C for 12 hours. The latter was distributed in 1 ml centrifuge tubes with 20% v/v glycerol and stored at

70°C, kept for later fermentations. The remaining 45 ml of initial culture were kept in

5 ml vials, by lyophilisation, using sterile skimmed milk as support at 10%

concentration and stored at 4°C.

b) Preparing pre-inoculums and inoculums

Pre-inoculums were prepared with the same medium composition corresponding to the batch; the microorganism conserved in solid sucrose medium was taken, then seeded in a volume of liquid medium, at 5-20% inoculum volume, cultured at 25-35 °C, with stirring at 100-400 rpm for 12-24 hours. Pre-inoculums were prepared with the same medium composition corresponding to the batch; the conserved microorganism was taken in solid sucrose medium, seeded in a volume of liquid medium, at 5-20% inoculum volume, cultured at 25-35 °C, with stirring at 100-400 rpm for 12-24 hours.

Composition of the medium used:

concentration (g/l)
10-20
0.03
0.02
0.002 - 0.1
0.0015 - 0.015
0.01-0.1

Carbon source:	
Sucrose	15-30
Nitrogen source:	
Yeast extract	15-30

The microorganism was seeded at 5-10 % of the fermentation volume and grown up to an average optical density of around 0.7 absorbance units in 1:10 dilution, read at 600nm. A sterile culture medium was used as target.

A preinoculum and inoculum must be made during fermentation, depending on fermenter volume, in such a way that the necessary quantity of cells is obtained in final inoculum (10% culture medium deposited in the production fermenter) to avoid the latency phase in the reactor and trying to maintain the 1:10 volume ratio between the preinoculum and the inoculum or sufficient cell density to serve as inoculum, maintaining rigorous control over culture purity and vegetative state so that it can be used as either inoculum or preinoculum.

c) Preparing the culture medium and inoculation

Culture medium pH was adjusted to pH 7.0. The balloon flask containing the medium for preparing the preinoculum was sterilised at 121°C for 15 minutes.

d) Operating conditions

Active ingredient was produced by batch fermentation using the established medium.

The operating conditions are listed in the following Table.

Fermenter operating conditions

Conditions	141
Medium volume (l)	10
Medium volume/fermenter volume ratio	0.8
Inoculum percentage	5-10
Inoculation optical density	0.5-1
Stirring (rpm)	100-400
Temperature (°C)	25-35
Aeration (vvm)	1-3
Initial medium pH	5-8
Fermentation time (hours)	6-12

2. Enzyme recovery:

a) Centrifuging

Extracellular enzyme was recovered by centrifuging at 5,000 rpm for 15 minutes for separating the biomass. The enzymatic extract or preparation presented 2-6 U/ml glucosyltransferase and fructosyltransferase activity. Extracellular enzyme was recovered by centrifuging at 5,000 rpm for 15 minutes to separate the biomass. The enzyme extract presented 2-6 U/ml glucosyltranspherase activity.

b) Ultrafiltration

Another way of recovering fermentation supernatant is by using 0.22-2 micra pore size ultra filtration membranes.

Another way of recovering the fermentation supernatant is by using 0.45-2 miera pore size ultra filtration membranes.

Example 3

Biopolymer production and recovery

a) Enzymatic reaction. Reaction conditions were as follows:

Reactant medium:

50-200 Mm phosphate buffer pH : 5 - 7

Substrate

: 8-20% sucrose

Enzyme quantity

: 10-30% v/v enzyme extract (200-500

U/D.

Reaction time

· 20-40 hours

Stirring

: 100-400 rpm

The enzyme was separated by centrifuging, placed in medium containing 8-20% sucrose, at pH 5-8 and 25-35°C for 20-30 hours, obtaining 30-60 g/l polymer concentration corresponding to 40-60% yield regarding substrate. Other processes reported to date have required up to 5-10 days for producing polymer. The reported microorganisms produced less polymer concentration (See Table 1). The enzyme was separated by centrifuging, placed in medium containing 8-20% sucrose, at pH 5-8 and at 25-35°C for 20-30 hours, obtaining 30-60 g/l polymer concentration corresponding to 40-60% yield regarding the substrate. In other reported processes 5-10 days for

producing the polymer were needed. The reported microorganisms produced less polymer concentration (Table 1).

b) Purifying the biopolymer

After the enzymatic reaction, the temperature was lowered to 4°C following enzyme reaction and it was possible to recover the biopolymer in two ways: The temperature was lowered to 4°C following the enzymatic reaction and it was possible to recover the biopolymer in two ways:

- Precipitation with solvents. 96% ethanol was added to cold reaction mixture with stirring. The quantity of added ethanol corresponded to 1.2-2.0 volumes of ethanol/volume reaction mixture.96% ethanol-was added to the cold reaction mixture with stirring. The quantity of added ethanol corresponded to 1.0-3.0 volumes of ethanol/volume of the reaction mixture.
- The precipitated biopolymer was dissolved in half the volume of deionised and distilled water and precipitated again with 1.2 to 2.0 volumes of ethanol/ reaction mixture volume. The precipitated biopolymer was redissolved in half the volume of deionised and distilled water and is precipitated again with 1.0 to 3.0 volumes of ethanol/ reaction mixture volume.
- Precipitated biopolymer was redissolved in a third of the volume of water and dried by lyophilisation or dried by compressed air at 60-80°C until reaching 5-10% humidity.

Table 1

EPS production using different microorganisms

Organism	Biopolymer (g/100 ml)	
	19 - 10 - 10 - 10 - 10 - 10 - 10 - 10 -	
Acetobacter pasteuria	inus	
ATCC 11142	0	
B. polymyxa		
NRRL B-68	0	
NRRL B-130	0	
NRRL B-510	1.2	
NRRL B-4317	1.4	
Isolate (NRRL B-184	75) 3.6	
B. subtilis		
NRRL B-447	1.0	
NRRL B-577	0	
NRRL B-644	0	
NRRL B-675	1.0	
NRRL B-744a	1.5	
NRRL B-2612	0	
Enterobacter levanicu	m	
NRRL B-1678	0.7	
Microbacterium laeva	niformans	
ATCC 15953	1.2	

<u>Ultrafiltration</u>. The reaction mixture was submitted to ultrafiltration on a regenerated cellulose membrane having a pore size greater than 10,000 Daltons to eliminate residual glucose and fructose. The biopolymer was then dryied by aspersion process.

Biopolymer production by this microorganism depends on the substrate concentration, this being optimal at 8-24% where the biopolymer is produced having the greatest degree of purity with the greatest yield (Table 2).

Table 2

Effect of sucrose on biopolymer production by Lactococcus lactis			
Sucrose (%) Biopolymer (g/l)		Biopolymer (g/l)	
-			
Control	0	0% (sucrose free)	
Sucrose	8	38.8	
Sucrose	12	50.1	
Sucrose	16	55.6	

c) Drying

The final product was obtained as a white powder that ean-which could be dried by lyophilisation or dry heat at a temperature not greater than 80°C. The obtained final product was a white powder which could be dried by lyophilisation or dry heat at a temperature not greater than 80°C.

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Example 4

Biopolymer characterisation

1. Solubility

The product was a hydro-soluble biopolymer able to form hydrogel homogeneous

dispersions up to 50% maximum concentration. 1.0 g of biopolymer was dissolved in

32 ml 5% chlorhydric acid, in 50 ml 10% sodium hydroxide or-and in 30 ml glacial

acetic acid.

It was insoluble in ethanol, isopropanol, acetone, mineral and vegetal oil and

polyethylen glycol.

The product was moderate soluble in 0.5% oxalic acid at ebullition temperature. The

product was moderately soluble in 0.5% oxalic acid at reflux temperature.

2. High Performance Liquid Chromatography (HPLC).

> A 1.5% biopolymer solution presented a 900-1,100 KDa molecular weight in

permeation chromatography determined on a Shodex OHPak KB-803 column.

Chromatography conditions were as follows:

Temperature : 55°C

Mobile phase : 0.1 M NaCl solution

Flow : 0.9 ml/min

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Polymer purity was greater than 95%, revealed by a thin peak in HPLC, in the following conditions:

Column: Shodex SC1011

Mobile phase: distilled deionised water

Flow: 0.6 ml/min.

Temperature: 70°C.

Equipment: Waters 510 with refraction index detector (Waters 2410).

The biopolymer presented a 7 to 7.5 minute retention time under these conditions.

The patterns used were analytic reagent grade glucose, fructose, and sucrose. The patterns used were analytic reagent grade glucose, fructose, sucrose and levan.

The biopolymer was stable over a broad range of pH shown by HPLC after contacting the polymer with pH 2-9 buffers. The biopolymer was stable over a broad pH range revealed by HPLC following polymer contact with pH 2-9 buffers.

3. Viscosity

Viscosity was determined in a 10% solution at 30°C using a ViscoEasy viscosimeter Serie L, Schott, Ref. 28.541.120, L2 stem at 50 rpm. The samples analysed presented viscosity ranging from 1,000-3,000 centipoises (cP). Pseudo-plastic behaviour was exhibited (cross-sectional thinning). Biopolymer solution viscosity became reduced on increasing the shear rate and increased on reducing temperature. Viscosity-was determined in a 10% solution at 30°C using a ViscoEasy viscosimeter L Series, Schott, Ref. 28.541.120, L2 stem at 50 rpm. The samples analysed presented viscosity ranging from 1,000-3,000 centipoises (eP). Pseudo plastic behaviour was exhibited (cross-sectional thinning). Biopolymer solution viscosity lowered whenthe shear rate increases and increases on reducing temperature.

4. Dimensional characteristics

The biopolymer had a true density close to that of sucrose (1.5 mg/ml). It is a material presenting high inter-particle porosity (48%). The biopolymer had a true density close to that of sucrose (1.5 mg/ml). It is a material showing high inter-particulate porosity (48%).

Average DVS particle size (diameter/volume/surface) was 224 micron.

5. Humidity adsorption

Water adsorption capacity ranged from 6.12 mg/g to 353.20 mg/g depending on relative humidity; this means that it was a slightly hygroscopic material. The biopolymer was capable of unlimited expansion on contact with water due to its polymeric structure and hydrophilicity, being able to form variable consistency systems depending on the quantity of water incorporated, giving rise to forming aqueous dispersions characterised by their high viscosity.

Water adsorption capacity ranged from 6.12 mg/g to 353,20 mg/g depending on relative humidity; this means that it was a slightly hygroscopic material. The biopolymer was able to soak up unlimited quantities of water due to its polymeric structure and hydrophilicity, being able to form variable consistency systems depending on the quantity of water incorporated, giving rise to the formation of aqueous dispersions characterised by their high viscosity.

6. Humidity

It presents losses of up to 10% when dried in a vacuum oven at 60°C.

7. Thermal characteristics

The biopolymer presentsed two vitreous transition points; the first between 20°C and 30°C and the second between 190°C and 220°C as determined by differential scanning calorimetry. The biopolymer presents two glass transition points; the first between 20°C and 30°C and the second between 190°C and 220°C as determined by scanning differential ealorimetry.

8. Microbiological quality

The biopolymer presents the following microbiological counts:

Microbiological charge	Range	<u>Unit</u>
Viable mesophile count	2000 - 4000	cfu/gr
Coliform count	Absence	mpn/gr
Faecal coliform count	≤10	mpn / gr
Salmonella count	Absence	
Mould and yeast count	<u>2000 - 5000</u>	cfu/gr

Microbiological charge	Range	Unit
Viable mesophile count	2000 4000	ufe / gr
Coliform count	Absence	nmp / gr
Faecal coliform count	<10	nmp / gr
Salmonella count	Absence	
mildew and yeast count	2000 5000	ufc / gr

9. Uses

- a) The biopolymer could be used in the pharmaceutical industry as viscosant, thickener, stabiliser, dispersant, as a film former, as disintegrant, blood plasma substitute, lubrication agent and/or prebiotic agent.
- b) The biopolymer could be used in the food industry as a thickener, viscosant, stabiliser, dispersant, as fibre and as fat , oils and ether- and ester-based carbohydrates substitute.
- c) The biopolymer can be used in products obtained by extrusion, for forming films apt for producing flexible and biodegradable packages and for obtaining disposable biodegradable products, obtained by injection or moulding and in the production of flocculent agents for water treatment.

ABSTRACT

A microorganism—was identified as a strain of Lactococcus lactis, NRRL B 30656, which when—cultivated and grown in a sucrose containing medium produces—a transferase extracellular—enzyme, which can be purified when is contacted with a sucrose—media, in suitable temperature and pH conditions, produces a glucose and fructose biopolymer

-A microorganism identified as a Lactococcus lactis strain (NRRL B-30656) produces an extracellular transferase enzyme when cultured and grown in sucrose-containing medium, which can be purified when it is brought into contact with a sucrose-based medium in suitable temperature and pH conditions, thereby producing a glucose and fructose polymer.